





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# Publisher Correction: Hypermethylation of gene body CpG islands predicts high dosage of functional oncogenes in liver cancer

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In the original version of this Article, the sixth sentence of the abstract incorrectly read ‘Most of the genes upregulated and with hypermethylated CGIs in the Alb-R26Met HCC model undergo the same change’, and should have read ‘Most of the genes upregulated and with hypermethylated CGIs in the Alb-R26Met HCC model undergo the same change in a large proportion of HCC patients’. This has been corrected in both the PDF and HTML versions of the Article.

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